Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).
The present invention relates to a device as defined by the preamble of claim 1.

A device of this kind is in frequent use in hospitals and is described, for example, in K. Seballe, “Migration of hydroxyapatite coated femoral prostheses”, Journal of Bone and Joint Surgery, volume 75-B, No. 5, September 1993, pp. 681-687.

US-A-5.577.089 discloses a method and equipment for measuring forms and orientations of bones in living beings. This patent gives an analysis of how e.g. the morphology of the vertebrae can be established based on bone density measurements by means of X-ray detection. Use is made of a computer which analyses data received and which uses the data to accurately define the shape and size of the vertebrae under investigation. Moreover, the computer is programmed to use the data to indicate the vertebral condition in medical terms.

US-A-5.577.089 describes measurements with respect to the human femur. Here, anatomically fixed points, like the proximal point and the medial epicondyle, are detected but they are only used to measure the femur length. Moreover, based on bone density measurements, a femur axis and a femur neck axis are calculated, as well as a femur head centerpoint. The patent discloses that these latter three features may be used to provide "an indication of any possible shifting of the prosthetic joint with respect to the femur" in case an artificial hip joint is implanted. Thus, US-A-5.577.089 discloses measuring shifting of a prosthetic joint relative to a bone supporting this joint, based on bone density measurements. However, the method proposed is very laborious since it needs the calculation of the intersection of two imaginary lines, and of an imaginary femur head centerpoint, for which many data elements of the femur and the prosthetic joint need be established.

WO-A-96/25086 is directed to providing a solution for the problem that prosthetic devices may be lost over time. To that end, this document discloses a method of evaluating bone density around a radiolucent composite prosthesis. Since the prosthetic device is transparent to X-rays the prosthesis is provided with three radioopaque reference markers embedded in the prosthesis. By means of a suitable densitometer, the boundaries of the prosthesis and the surrounding bone are identified and stored for later use. A region of interest is defined which is the area of the bone adjacent to the prosthesis. In this region of interest, the density of the bone is measured over time in order to establish any degradation of the bone to which the prosthesis is fixed. From the measurement data, the loosening of the prosthesis from the bone can be established.

The locations of the three radio-opaque markers are also stored in order to compare measurements later in time with prior measurements. To this end, a stored template with three template reference markers is used which are fitted to measured reference markers. To be sure that later measurements can be compared with former measurements, also, in the case of a hip implant, also the location of the lesser trochanter is identified and stored. Thus, both the three reference markers and the lesser trochanter are used to ensure that that subsequent scans of a patient will be properly aligned and may be used for direct comparison with earlier scans.

A device defined by the preamble of claim 1 is used, for example, for determining the position of a hip prosthesis with respect to the femur, to which the hip prosthesis is connected on one side. On the other side, the hip prosthesis is in contact with an acetabular prosthesis, which is connected to the pelvis.

Wear to the hip joint leads to a very painful limitation of the movements which a person is able to carry out. Since the 1970s, hip prostheses have been widely used in orthopaedics to replace a hip joint which has become worn. However, treatment of the arthrosis does not end with the fitting of a hip prosthesis, but rather in practice is the beginning of a long period of careful monitoring of the patient. The monitoring consists both of physical examinations and of the study of regular X-rays.

When the method was introduced, the minimum age of patients to be treated was approximately 70 years. However, nowadays hip prostheses of this kind are also fitted to people of an increasingly young age. Younger people have a higher activity level than older people, with the result that hospitals are confronted to an increasing extent with hip prostheses which become detached from the bone to which they are connected.

Figure 1 diagrammatically shows an X-ray of a hip prosthesis, which at the top is in contact, by means of a spherical end, with an acetabular prosthesis which is attached to the pelvis, and at the bottom is connected by means of a pin to the femur.

Just below the hip joint, the femur has two marked projections, the outer projection being referred to as the trochanter major and the inner projection as the trochanter minor. The surface of the trochanter major is rough, so as to increase the contact area for the attached gluteus and thigh muscles. The trochanter minor lies on the inside and points 30° towards the rear. Only one muscle is attached to the trochanter minor, and this muscle, when tightened, causes the hip joint to bend and the femur to rotate outwards. Both tubercles are situated at a fixed location. This means that the shape, the location with respect to the leg and the size are not affected by positioning a prosthesis in the femur. The trochanter major, the trochanter minor, as well as the axis of the knee joint (not shown in Figure 1), are situated at fixed anatomical positions which form orientation points for the correct positioning of the prosthesis.

If a hip prosthesis becomes detached from the femur, the result is that the prosthesis slowly sinks into the femur, causing damage to the femur. If such an event is only discovered at a late stage, considerable amounts of bone may already have been lost, and this
first has to be replaced with donor bone in order to repair the anatomy to a sufficient extent for the same prosthesis 1 to be replaced.

[0013] The "moment" at which the mechanical detachment occurs is not precisely known. With standard current X-ray techniques, it is only possible to detect whether a prosthesis is attached or has become detached, or at least whether the prosthesis 1 has moved more than 5 mm. In medical circles, the assumption is that the increase in the speed of migration is the "moment" of detachment. The speed of migration is understood to mean the rate at which the prosthesis 1 moves with respect to the femur 6.

[0014] The above-mentioned article by Søballe describes a standard procedure which can be used to measure the current position of the prosthesis 1 with respect to the femur 6. This method is known as the X-ray stereophotogrammetry analysis (RSA).

[0015] This standard procedure can be used to measure the movement of the prosthesis 1 in the femur 6 to an accuracy of 0.1 mm. In order to be able to make use of this standard procedure, during the hip operation it is necessary to arrange various small tantalum balls, which usually have a diameter of 0.8 mm, at various locations in the bone before fitting the hip prosthesis 1. Moreover, the prosthesis 1 itself also has to be provided with at least one small tantalum ball, which serves as a reference location. The small tantalum balls arranged in the bone no longer move after the hip operation.

[0016] In the RSA procedure, two X-ray cameras 10, 11 are used to take at least two different X-rays from different directions. The X-radiation is directed in such a way that the small tantalum balls in the femur 6 and on the prosthesis 1 are visible. By making use of the two pictures, which are taken from different angles, and a known trigonometric measurement, the spatial position of the prosthesis 1 can be accurately determined with respect to the femur 6. By repeating such measurements over the course of time, the migration of the prosthesis 1 with respect to the femur 6 can be determined. Measuring the migration of an implant relative to a bone to which the implant is connected, especially during the first year after the implant has been implanted, appears to be a good indication for possible future mechanical loosening, as is also indicated by another document directed to the RSA method: L. Ryd, Roentgen Stereophotogrammetric Analysis of Prosthetic Fixation in the Hip and Knee Joint, Clinical orthopaedics and Related Research, Number 276, March, 1992.

[0017] Although the above-mentioned RSA procedure is extremely accurate, it is also extremely laborious. Moreover, this known method can only be used on a select group of patients, since only a few teaching hospitals have the advanced equipment which is required.

[0018] It is therefore desirable to provide a device which can be used to determine the migration of an implant, which is connected to a bone, in a body with a high level of accuracy but without extra actions, such as the attachment of small tantalum balls, being required during the operation prior to the attachment of the implant.

[0019] To achieve this object, the device of the above-mentioned type is improved as defined in claim 1.

[0020] As is evident from a.o. US-A-5,777,089 modern shape recognition means are available with which locations of anatomically fixed points on bones can be established. According to the invention, the locations of one or more of such anatomically fixed points is used to establish the relative displacement between bones and implants connected to the bones. The method which is used in the device according to the invention is straightforward and uses only a limited number of process steps. No calculation of imaginary lines and centers is required to achieve a very reliable result.

[0021] The device according to the invention is not only applicable to implants portions of which are inserted into bone portion but also to medical supporting structures connected to the outside of bones. Therefore, for the purpose of this invention, "implants" are defined to include such supporting structures. It is no longer necessary to use, for example, small tantalum balls, the position of which is established with the aid of X-rays, but rather it is sufficient to use means for establishing the position of the at least one anatomically fixed point with respect to the prosthesis. In this case, to establish the position of the prosthesis, use is made of the location of, e.g., two identifying marks which are connected to the prosthesis. These identifying marks may, for example, as in the prior art, comprise small objects which can be located with the aid of X-rays, for example small tantalum balls. However, since it is nowadays possible to detect accurately shapes of objects, it is also possible to select, preferably, two fixed points on the prosthesis itself to be identifying marks, the location of which is established with the aid of shape recognition means. In this case too, there are therefore three known locations, with the aid of which respective positions can be determined with the aid of trigonometry.

[0022] It is not necessary for the method to make use of, e.g., two objects or points, which are to be located, on the implant and at least one bone marking. For example, one can alternatively use two anatomically fixed points on the bone and one identifying mark on the implant.

[0023] In a first preferred embodiment the generator means of the device comprises

A1. first generating means for generating first radiation at a first position and directing the first radiation onto the at least one bone marking (7, 8; 13, 14) and the at least one predetermined identifying mark (4, 5) of the implant (1; 15, 15') from a first direction;

A2. second generating means for generating second radiation at a second position and directing the second radiation onto the at least one bone marking (7, 8; 13, 14) and the at least one predetermined identifying mark (4, 5) of the implant (1; 15, 15') from a second direction.
second direction;
and said receiving means are arranged for:

B1. receiving first and second radiation images, respectively, of said at least one bone marking and said at least one predetermined identifying mark, formed by said first and second radiation, respectively;

and said evaluation means are arranged for:

C1. determining the position of the implant (1; 15, 15') with respect to the bone (6; 16) on the basis of said first and second radiation images received by said receiving means. The implant may, for example, be a hip prosthesis, in which case the bone marking may be selected from the following two anatomically fixed points: the trochanter major and the trochanter minor.

[0024] However, the device may also be used to determine migration of a knee prostheses, in which case the bone marking is selected from the following two anatomically fixed points: the medial epicondyle and the lateral epicondyle.

[0025] Furthermore, the device according to the present invention may be used for any other form of prosthesis which is positioned in a bone where the bone has clearly recognizable, anatomically fixed points. The device is advantageous above all (but not exclusively) in connection with joint-replacement implants, since all joints have unique bone markings.

[0026] The invention will be explained in more detail below with reference to several drawings, which are intended only to illustrate the invention and not to limit it. In the drawings:

Figure 1 shows a diagrammatic picture of a hip prosthesis which is connected to the femur;
Figure 2 diagrammatically shows a knee prosthesis which is connected to the femur and the tibia.

[0027] The elements associated with the reference numerals 1 to 11 have already been described above. The reference numeral 12 indicates an evaluation device, for example a computer, which is connected to the radiation sources 10, 11. The radiation sources 10, 11 are designed to generate X-radiation. However, it is also conceivable that radiation of a different frequency may be generated and used for the purposes of the present invention. Theoretically, even ultrasound sources may be used.

[0028] The evaluation means 12 are designed, inter alia, to control the radiation sources 10, 11.

[0029] Furthermore, two receivers 17, 18 are provided for receiving the radiation emitted by the radiation sources 10, 11, after the radiation has radiated through the prosthesis with the surrounding bone. The receivers 17, 18 are connected to the evaluation device 12 for the purpose of transmitting the images which they receive.

[0030] The evaluation means 12 preferably comprise a computer, the memory of which has been loaded with a software program for recognizing shapes of bones. A program which can advantageously be used is the Scipio™ program, which has already been in use for some time in bone banks. Bone banks are establishments where bones are stored for subsequent use in transplants. The shape of bones stored in bone banks can be recognized with the aid of CCDs and the Scipio™ program. The shapes of the bones stored are recorded and held in a memory of a computer. In the event of requests for bones to be supplied, shapes of bones which have been requested can be compared with recorded shapes of bones, so that bones for transplant purposes can be supplied more easily and more quickly.

[0031] The above-mentioned Scipio™ program is in principle able to locate anatomically fixed points on bones.

[0032] The above-mentioned Scipio™ program can therefore in principle be used, for example, to locate the trochanter minor 7 and the trochanter major 8 of the femur. These points can then be used instead of the locations of the small tantalum balls which in the prior art are placed in the bone in order to determine the position of the femur 6.

[0033] To determine the position of the hip prosthesis 1, use can be made of the small tantalum balls 4, 5 which are connected to the prosthesis 1. However, there is no need to use these small tantalum balls. It will be clear that the Scipio™ program can recognize and record not only the shapes of bones, but also the shapes of implants. The Scipio™ program can therefore in principle also record and locate fixed points on implants. Hence it is then possible to carry out a trigonometric measurement which is known per se in order to establish the respective positions of the implant 1 and the bone 6.

[0034] It will be clear that the trigonometrical measurement only needs three points to be located. To do this, it is in principle unimportant whether two of the three points are connected to the bone 6 and one to the implant 1 or, as an alternative, one point is connected to the implant 1 and two of the three points are connected to the bone. It is generally quite possible to locate two anatomically fixed points on the bone. As has been mentioned, the femur 6 has two anatomically fixed points, namely the trochanter minor 7 and the trochanter major 8.

[0035] Figure 2 diagrammatically shows a knee prosthesis 15, 15', which is connected to the femur 6 and the tibia 16.

[0036] In this case too, it is possible to make use of one or more anatomically fixed points, namely the medial epicondyle 13 and the lateral epicondyle 14.

[0037] Here too, it is the case that use may be made of small tantalum balls 4, 5 or that use may be made of a program which can record the shape of the knee pros-
thesis 15, 15' and then locate one or two fixed points thereon.

[0038] It will be clear that instead of trigonometrical measurements it is also in principle possible to use measurements which use more points.

[0039] The reference numerals 10, 11, 12, 17 and 18 in Figure 2 refer to the same components as in Figure 1.

[0040] Since the device described above no longer make use of separate small tantalum balls or the like placed in the bone, the above-mentioned method can easily be employed in all hospitals where prostheses are fitted.

[0041] This means that the migration of a prosthesis with respect to the surrounding bone can be established in a simple and rapid manner.

[0042] It is extremely important that the migration of a prosthesis with respect to the surrounding bone is determined above all in the first few months after the prosthesis is fitted. This is because the extent of migration in the first few months has been found to be a measure of the probability of the prosthesis becoming detached. If it is found that the migration in the first few months is greater than a defined threshold, it can be decided to perform a surgical intervention, which can prevent needless disintegration of the bone. This means that interventions can be carried out more quickly, and there is less need to carry out revision operations in which damaged bone first has to be replaced before a new prosthesis can be fitted. This reduces the operation time and therefore saves considerable expense for the health service.

[0043] The receivers 17, 18 can be standard commercially available receivers which are provided with an image intensifier which is known per se. For this reason, there is no need for any more radiation for taking these X-rays, for example, than when taking conventional X-rays.

Claims

1. Device for measuring the position of an implant (1; 15, 15') relative to at least one bone (6; 16) in a body, to which bone the implant is connected, which bone (6; 16) has at least one bone marking (7, 8; 13, 14), and which implant has at least one predetermined identifying mark (4, 5), provided with the following means:

   A. generator means (10, 11) for generating radiation and directing said radiation onto said at least one bone marking and said at least one predetermined identifying mark;

   B. receiving means (17, 18) for receiving a radiation image of said at least one bone marking and said at least one predetermined identifying mark;

   C. evaluation means (12), which are coupled to the receiving means (17, 18), and adapted for determining the position of the implant (1; 15, 15') with respect to the bone (6; 16) on the basis of the radiation image received during operation of means B;

   D. means for repeating the operation of means A, B and C over time to determine migration of the implant with respect to said at least one bone;

characterized in that the evaluation means are provided with means adapted for recognizing the shape of the at least one bone (6; 16), for locating the at least one bone marking on the basis of an anatomically fixed point (7, 8; 13, 14) on the bone (6; 16) and said evaluation means are arranged for establishing the position of said at least one anatomically fixed point with respect to said at least one predetermined identifying mark of said implant.

2. Device according to claim 1, wherein said generator means comprise:

   A1. first generating means (10) for generating first radiation at a first position and directing the first radiation onto the at least one bone marking (7, 8; 13, 14) and the at least one predetermined identifying mark (4, 5) of the implant (1; 15, 15') from a first direction;

   A2. second generating means (11) for generating second radiation at a second position and directing the second radiation onto the at least one bone marking (7, 8; 13, 14) and the at least one predetermined identifying mark (4, 5) of the implant (1; 15, 15') from a second direction;

   and said receiving means (17, 18) are arranged for:

   B1. receiving first and second radiation images, respectively, of said at least one bone marking and said at least one predetermined identifying mark, formed by said first and second radiation, respectively;

   and said evaluation means (12) are arranged for:

   C1. determining the position of the implant (1; 15, 15') with respect to the bone (6; 16) on the basis of said first and second radiation images received by said receiving means.

3. Device according to claim 1 or 2, wherein said evaluation means are arranged for determining said position on the basis of said radiation image comprising at least two predetermined identifying marks (4, 5) on the implant.

4. Device according to any of the claims 1-3, wherein said evaluation means are arranged for determining
said position on the basis of said radiation image comprising at least two anatomically fixed points (7, 8; 13, 14) on the bone (6; 16).

5. Device according to any of the claims 1-4, wherein the implant is a hip prosthesis (1), the generator means are X-radiation generator means and the predetermined identifying marks of the implant are objects (4, 5) which are impermeable to X-radiation.

6. Device according to claim 5, wherein the evaluation means (12) are arranged to recognize either the trochanter major or trochanter minor, or both, as the at least one bone marking.

7. Device according to any of the claims 1 through 4, wherein the implant is a knee prosthesis (15, 15'), the generator means are X-radiation generator means and the predetermined identifying marks of the implant are objects (4, 5) which are impermeable to X-radiation.

8. Device according to claim 7, wherein the evaluation means (12) are arranged to recognize either the medial epicondyle or lateral epicondyle, or both, as the at least one bone marking.

Patentansprüche

1. Vorrichtung zum Messen der Position eines Implantats (1; 15, 15') relativ zu mindestens einem Knochen (6; 16) in einem Körper, wobei das Implantat mit dem Knochen verbunden ist, wobei der Knochen (6; 16) mindestens eine Knochenmarkierung (7, 8; 13, 14) hat und wobei das Implantat mindestens eine vorbestimmte Identifizierungsmarkierung (4, 5) hat, wobei die Vorrichtung mit den folgenden Einrichtungen versehen ist:

   A. Erzeugungseinrichtungen (10, 11) zur Erzeugung von Strahlung und zum Ausrichten der Strahlung auf die mindestens eine Knochenmarkierung und die mindestens eine vorbestimmte Identifizierungsmarkierung;
   B. Empfangseinrichtungen (17, 18) zum Empfangen eines Strahlungsbilds der mindestens einen Knochenmarkierung und der mindestens einen vorbestimmten Identifizierungsmarkierung;
   C. Beurteilungseinrichtungen (12), die mit den Empfangseinrichtungen (17, 18) gekoppelt sind, die Position des Implantats (1; 15, 15') zu bestimmen;
   D. Einrichtungen zum Wiederholen des Betriebs der Einrichtungen A, B und C über die Zeit, um eine Migration des Implantats in bezug auf den mindestens einen Knochen zu bestimmen;

dadurch gekennzeichnet.

2. Vorrichtung nach Anspruch 1, wobei die Erzeugungseinrichtungen folgendes aufweisen:

   A1. eine erste Erzeugungseinrichtung (10) zum Erzeugen einer ersten Strahlung in einer ersten Position und zum Ausrichten der ersten Strahlung auf die mindestens eine Knochenmarkierung (7, 8; 13, 14) und die mindestens eine vorbestimmte Identifizierungsmarkierung (4, 5) des Implantats (1; 15, 15') aus einer ersten Richtung;
   A2. eine zweite Erzeugungseinrichtung (11) zum Erzeugen einer zweiten Strahlung in einer zweiten Position und zum Ausrichten der zweiten Strahlung auf die mindestens eine Knochenmarkierung (7, 8; 13, 14) und die mindestens eine vorbestimmte Identifizierungsmarkierung (4, 5) des Implantats (1; 15, 15') aus einer zweiten Richtung;
   B1. erste bzw. ein zweite Strahlungsbilder der mindestens einen Knochenmarkierung und der mindestens einen vorbestimmten Identifizierungsmarkierung zu empfangen, die von der ersten bzw. zweiten Strahlung gebildet sind;
   C1. die Position des Implantats (1; 15, 15') in bezug auf den Knochen (6; 16) auf der Basis der von den Empfangseinrichtungen empfangenen ersten und zweiten Strahlungsbilder zu bestimmen.

3. Vorrichtung nach Anspruch 1 oder 2, wobei die Beurteilungseinrichtungen angeordnet sind, um:

   B1. erste bzw. ein zweite Strahlungsbilder der mindestens einen Knochenmarkierung und der mindestens einen vorbestimmten Identifizierungsmarkierung zu empfangen, die von der ersten bzw. zweiten Strahlung gebildet sind;
   C1. die Position des Implantats (1; 15, 15') in bezug auf den Knochen (6; 16) auf der Basis der von den Empfangseinrichtungen empfangenen ersten und zweiten Strahlungsbilder zu bestimmen.
stimmte Identifizierungsmarkierungen (4, 5) an dem Implantat aufweist.

4. Vorrichtung nach einem der Ansprüche 1 bis 3, wobei die Beurteilungseinrichtungen angeordnet sind, um die Position auf der Basis des Strahlungsbilds zu bestimmen, das mindestens zwei anatomiche Fixpunkte (7, 8; 13, 14) an dem Knochen (6; 16) aufweist.

5. Vorrichtung nach einem der Ansprüche 1 bis 4, wobei das Implantat eine Hüftgelenkprothese (1) ist, wobei die Erzeugungseinrichtungen Röntgenstrahlungs-Erzeugungseinrichtungen sind und wobei die vorbestimmten Identifizierungsmarkierungen des Implantats Objekte (4, 5) sind, die für Röntgenstrahlung undurchlässig sind.

6. Vorrichtung nach Anspruch 5, wobei die Beurteilungseinrichtungen angeordnet sind, um entweder den Trochanter major oder den Trochanter minor oder beide als die mindestens eine Knochenmarkierung zu erkennen.

7. Vorrichtung nach einem der Ansprüche 1 bis 4, wobei das Implantat eine Kniegelenkprothese (15, 15') ist, wobei die Erzeugungseinrichtungen Röntgenstrahlungs-Erzeugungseinrichtungen sind und wobei die vorbestimmten Identifizierungsmarkierungen des Implantats Objekte (4, 5) sind, die für Röntgenstrahlung undurchlässig sind.

8. Vorrichtung nach Anspruch 7, wobei die Beurteilungseinrichtungen (12) angeordnet sind, um entweder den Epicondylus medialis oder den Epicondylus lateralis oder beide als die mindestens eine Knochenmarkierung zu erkennen.

Revendications

1. Dispositif pour mesurer la position d’un implant (1 ; 15, 15') par rapport à au moins un os (6 ; 16) dans un corps, os auquel l’implant est relié, cet os (6 ; 16) comportant au moins un marquage d’os (7, 8 ; 13, 14), et cet implant comportant au moins une marque d’identification prédéterminée (4, 5), muni des moyens suivants :

A. des moyens formant générateur (10, 11) pour générer un rayonnement et diriger ledit rayonnement sur ledit marquage d’os au nombre d’au moins un et ladite marque d’identification prédéterminée au nombre d’au moins une ;

B. des moyens de réception (17, 18) pour recevoir une image de rayonnement dudit marquage d’os au nombre d’au moins un et de ladite marque d’identification prédéterminée au nombre d’au moins une ;

C. des moyens d’évaluation (12), qui sont couplés aux moyens de réception (17, 18), et adaptés pour déterminer la position de l’implant (1 ; 15, 15') par rapport à l’os (6 ; 16) en fonction de l’image de rayonnement reçue durant le fonctionnement des moyens B ;

D. des moyens pour répéter le fonctionnement des moyens A, B et C au cours du temps pour déterminer la migration de l’implant par rapport audit os au nombre d’au moins un ;

4. Dispositif selon la revendication 1, dans lequel lesdits moyens formant générateur comprennent :

A1. des premiers moyens de génération (10) pour générer un premier rayonnement en une première position et diriger le premier rayonnement sur le marquage d’os au nombre d’au moins un (7, 8 ; 13, 14) et la marque d’identification prédéterminée au nombre d’au moins une (4, 5) de l’implant (1 ; 15, 15') à partir d’une première direction ;

A2. des deuxième moyens de génération (11) pour générer un deuxième rayonnement en une deuxième position et diriger le deuxième rayonnement sur le marquage d’os au nombre d’au moins un (7, 8 ; 13, 14) et la marque d’identification prédéterminée au nombre d’au moins une (4, 5) de l’implant (1 ; 15, 15') à partir d’une deuxième direction ;

et lesdits moyens de réception (17, 18) sont configurés pour :

B1. recevoir des première et deuxième images de rayonnement, respectivement, dudit marquage d’os au nombre d’au moins un et de ladite marque d’identification prédéterminée au nombre d’au moins une, formées par lesdits premiers et deuxième rayonnements, respectivement ;

et lesdits moyens d’évaluation (12) sont configurés pour :

C1. déterminer la position de l’implant (1 ; 15,
3. Dispositif selon la revendication 1 ou 2, dans lequel lesdits moyens d'évaluation sont configurés pour déterminer ladite position en fonction de ladite image de rayonnement comprenant au moins deux marques d'identification prédéterminées (4, 5) sur l'implant.

4. Dispositif selon l'une quelconque des revendications 1 à 3, dans lequel lesdits moyens d'évaluation sont configurés pour déterminer ladite position en fonction de ladite image de rayonnement comprenant au moins deux points anatomiquement fixes (7, 8 ; 13, 14) sur l'os (6 ; 16).

5. Dispositif selon l'une quelconque des revendications 1 à 4, dans lequel l'implant est une prothèse de la hanche (1), les moyens formant générateur sont des moyens formant générateur de rayons X et les marques d'identification prédéterminées sur l'implant sont des objets (4, 5) qui sont imperméables aux rayons X.

6. Dispositif dont la revendication 5, dans lequel les moyens d'évaluation (12) sont configurés de façon à reconnaître soit le grand trochanter soit le petit trochanter, ou les deux, comme étant le marquage d'os au nombre d'au moins un.

7. Dispositif selon l'une quelconque des revendications 1 à 4, dans lequel l'implant est une prothèse du genou (15, 15'), les moyens formant générateur sont des moyens formant générateur de rayons X et les marques d'identification prédéterminées de l'implant sont des objets (4, 5) qui sont imperméables aux rayons X.

8. Dispositif selon la revendication 7, dans lequel les moyens d'évaluation (12) sont configurés de façon à reconnaître soit l'épicondyle médian soit l'épicondyle latéral, ou les deux, comme étant le marquage d'os au nombre d'au moins un.
REFERENCES CITED IN THE DESCRIPTION

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Patent documents cited in the description

• US 5577089 A [0003] [0004] [0004]
• WO 9625086 A [0005]

Non-patent literature cited in the description